Genome analysis

# Data structures and compression algorithms for genomic sequence data

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#### **ABSTRACT**

Motivation: The continuing exponential accumulation of full genome data, including full diploid human genomes, creates new challenges not only for understanding genomic structure, function and evolution, but also for the storage, navigation and privacy of genomic data. Here, we develop data structures and algorithms for the efficient storage of genomic and other sequence data that may also facilitate querying and protecting the data.

Results: The general idea is to encode only the differences between a genome sequence and a reference sequence, using absolute or relative coordinates for the location of the differences. These locations and the corresponding differential variants can be encoded into binary strings using various entropy coding methods, from fixed codes such as Golomb and Elias codes, to variables codes, such as Huffman codes. We demonstrate the approach and various tradeoffs using highly variables human mitochondrial genome sequences as a testbed. With only a partial level of optimization, 3615 genome sequences occupying 56 MB in GenBank are compressed down to only 167 KB, achieving a 345-fold compression rate, using the revised Cambridge Reference Sequence as the reference sequence. Using the consensus sequence as the reference sequence, the data can be stored using only 133 KB, corresponding to a 433-fold level of compression, roughly a 23% improvement. Extensions to nuclear genomes and high-throughput sequencing data are discussed.

Availability: Data are publicly available from GenBank, the HapMap web site, and the MITOMAP database. Supplementary materials with additional results, statistics, and software implementations are available from http://mammag.web.uci.edu/bin/view/Mitowiki/ ProjectDNACompression.

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## 1 INTRODUCTION

As high-throughput genome sequencing technologies continue to improve, genome sequence data continue to accumulate at an exponential pace. Not only do we already have the genome sequence of thousands of viruses and bacteria and dozens of multicellular organisms from plants to humans, but we are rapidly approaching the stage where sequencing individual diploid human genomes will be economically affordable. The first diploid human genome sequences were recently produced (Levy et al., 2007; Wang et al., 2008; Wheeler et al., 2008) and a project to sequence 1000 human genomes in the next few years is under way (Kaiser, 2008). And so is the race for the capability to sequence an individual human genome for less than \$1000 within a few years (Service, 2006). Millions of human genome sequences could be generated within a decade or two.

In addition to the obvious challenges to understand the structure, function and evolution of genomes, modern high-throughput sequencing (HTS) methods also raise questions about how to efficiently represent, store, transmit, query and protect the privacy of sequence information. These questions are further reinforced if one takes into account also progress in synthetic biology and our ability to bioengineer new sequences.

Currently, publicly available genomes are typically stored as flat text files in GenBank, but this approach is unlikely to scale up in many ways. The storage of the diploid genomes of all currently living humans using this simple approach would take 'GenBank', without counting headers or any additional annotations, on the order of  $36 \times 10^{18}$  bytes, or 36M Terabytes, an amount difficult to store or download over the Internet, even using standard compression technologies (e.g. gzip). And even with the progress that can be expected with Moore's law for storage and networking in the coming years, it is likely that security and privacy issues will require additional layers of protection around genomic data.

Here, we develop data structures and algorithms to begin addressing these problems. These data structures allow the compression of genome and other sequences while facilitating certain classes of sequence queries by bypassing classical sequence alignments and dynamic programming algorithms. The approach is demonstrated primarily using a benchmark dataset comprising a few thousand individual mitochondrial genomes sequences. Human mitochondrial sequences provide an excellent testbed for developing and testing efficient data structures and algorithms because, unlike nuclear genome sequences, many thousands of fully sequenced mitochondrial genomes are already available, from a diverse population of individuals. In addition, mitochondrial genome sequences pose unique challenges due to their greater variability, as compared with single nucleotide polymorphism (SNP) data.

#### 2 GENERAL APPROACH

In the case of multiple genomes from the same species, associated with 'resequencing' technologies, the flat text file approach is clearly wasteful since for the most part the sequences are identical. Thus a

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simple approach is to store a reference sequence, and then for each other sequence, encode only the differences (or 'deltas') with respect to the original sequence. More precisely, consider first the sequences AACGACTAGTAATTTG and CACGTCTAGTAATGTG which are identical, except for a substitution in position 1 (A $\rightarrow$ C), 5 (A $\rightarrow$ T) and 14 (T $\rightarrow$ G). Each SNP can be encoded by a pair (i,X), where i is an integer encoding the position and X represents the value of the substitution relative to the reference. Thus given the first sequence as a reference, the second one can be encoded by the string '1C5T14G', concatenating the coordinates of the locations at which the variations occur and the SNP values at these locations. Note that with this data representation, the questions 'Is this sequence different from the reference sequence at position i? And if so how?' are easy to answer. Thus, the same data structure that facilitates compact representation, facilitates also efficient information retrieval.

Other events such as deletions and insertions can easily be accommodated in the same general scheme. For a deletion, imagine using two integers (i,l) where the first integer denotes the position where the deletion occurs, and the second integer represents the length of the deletion. Likewise, for an insertion of length l, one can use the encoding  $i, X_1 ... X_l$  to denote the insertion of  $X_1 ... X_l$  at position i with respect to the reference sequence.

Although the basic idea is easy to understand, and not new, a precise implementation requires addressing a number of important technical issues. A first observation is that one can use local relative addresses, i.e. intervals, rather than absolute addresses. Using intervals, the above example '1C5T14G' becomes '0C4T9G'. With intervals the dynamic range of the integers to be encoded may be considerably smaller than with absolute addresses. The relatively modest price to pay is that intervals must be added to recover absolute coordinates.

A second observation is that if the positions at which variations occur in the population are fixed and form a relatively small subset of all possible positions, then additional savings may result by focusing only on those positions. If in the same schematic example as above, one knew that the population substitutions can occur only at positions 1, 5 and 14, then one could, for instance, encode '1C5T14G' simply by 'CTG', at the cost of keeping an additional table storing the coordinates where the variants occur, and using the letter in the reference sequences at positions where the reference sequence and the sequence under consideration are identical. This approach could be suitable, for instance, for the SNP HapMap data (The International HapMap Consortium, 2003, 2007), but may not be suitable in other situations, where either the location of all possible variations occurring in the population under consideration is not known in advance, or the number of such locations is very large across the population, but not very large in a typical sequence. This is the case, for instance, of mitochondrial DNA which is characterized by much higher mutation rates than nuclear DNA. Thus, different situations may lead to different variations of the basic

An additional technical consideration is the choice of the reference sequence. In particular, the reference sequence does not need to be an actual genome but can, for instance, correspond to a consensus genome. While the resequencing case is of primary interest here due to the medical implication associated with resequencing human genomes, the same general ideas can be applied also to the case of *de novo* sequencing by using, for instance, the genome of the closest available species as the reference genome.

However, no matter what the detailed scenario is, all applications of the basic ideas hinge on a fundamental technical problem: how to encode integers, representing for instance absolute or relative genomic addresses or read lengths, into binary strings. It is essential to understand that the naive idea of converting integers to their binary value, that is, converting a '5' to '101' does not work at all since with this encoding one does not know where an integer ends and the next one begins. There are no spaces, tabs or commas available to separate consecutive integers in the ultimate binary format of any computer where only the symbols 0 and 1 are available. Thus, the encoding itself must somehow contain the information necessary to uniquely determine the beginning and end of each information item. In addition, the plain conversion of integers to binary does not take into account any entropy considerations. Similarly, a general purpose compression scheme for text data, such as Lempel-Ziv (gzip), is likely to be far from optimal for genome and HTS data. In short, we are interested in binary encoding schemes for sequences of integers that can be parsed automatically and that, consistently with information theory, are entropy efficient, in the sense that fewer bits are used to encode more frequent events. The goal here is not to prescribe a single strategy to achieve this end, but rather to present a family of related coding strategies and some of the tradeoffs that would have to be optimized in a practical application, and illustrate the approach using highly variable mitochondrial DNA.

#### 3 SPECIFIC ENCODING STRATEGIES

To begin with, we illustrate these issues here by considering how the integer positions i are ultimately encoded into a binary string. From Shannon's entropy coding theory (Cover and Thomas, 1991; McEliece, 1977), optimal encoding of these integers from a compression standpoint depends on their distribution in order to assign shorter binary codes to more probable symbols (integers). For simplicity, we distinguish two broad classes of codes: fixed codes, such as Golomb codes (Golomb, 1965) and Elias codes (Elias, 1975), and variable codes, such as Huffman codes (Huffman, 1952). In a fixed code, the integer i is always encoded in the same way, whereas in a variable code the encoding changes.

# 3.1 Fixed codes: Golomb and Golomb-Rice codes

Both Golomb codes and Elias codes encode an integer j by concatenating two bit strings: a preamble p(j), that encodes j's scale, and a mantissa. Golomb codes were specifically developed to encode stationary coin flips with  $p \neq 0.5$ . Thus, they are known to be optimal and asymptotically approach the Shannon limit if the data are generated by random coin flips or, equivalently, if the distribution over the integers is geometric, although they can be used for any other distribution. The more skewed the probability p is (towards 0 or 1) the greater the level of compression that can be achieved.

Golomb codes have one integer parameter m. Given m, any positive integer j can be written using its quotient and remainder modulo m as  $j = \lfloor j/m \rfloor + (j \mod m)$ . To encode j, the Golomb code with parameter m (Table 1) encodes the quotient and remainder by using:

- $\lfloor j/m \rfloor$  1-bits for the quotient;
- followed by a 0, as a delimiter (unary encoding of |j/m|);

**Table 1.** Golomb encoding of the integers j = 0 to 8, for different values of the parameter m

j	m=2	m=3	m=4	m=5	m=6
0	00	00	000	000	000
1	01	010	001	001	001
2	100	011	010	010	0100
3	101	100	011	0110	0101
4	1100	1010	1000	0111	0110
5	1101	1011	1001	1000	0111
6	11100	1100	1010	1001	1000
7	11101	11010	1011	1010	1001
8	111100	11011	11000	10110	10100

 followed by the phased-in binary code for j mod m for the remainder (described below).

The encoding of integers 0, ..., m-1 normally requires  $B = \lceil \log m \rceil$  bits. If m is not a power of two, then one can sometimes use B-1 bits. More specifically, in the 'phased-in' approach:

- if  $i < 2^B m$ , then encode i in binary, using (B-1) bits;
- if  $i \ge 2^B m$ , then encode i by  $i + 2^B m$  in binary, using B bits.

For instance, for m=5, i=2 is encoded as '10' using 2 (=B-1) bits, and i=4 is encode as '111' using 3 (=B) bits (Table 1). Thus the encoding of j requires in total  $\lfloor j/m \rfloor + 1 + \lfloor \log m \rfloor$  or  $\lfloor j/m \rfloor + 1 + \lceil \log m \rceil$  bits (Table 1) and the codeword for the integer j+m has one more bit than the codeword for the integer j. Unless otherwise specified, all logarithms are taken to base 2. We use also ' $\lceil \log m \rceil$ ' to denote ' $\lceil \log m \rceil$  or  $\lceil \log m \rceil$ '.

The entropy of the geometric distribution of the coin flip runlengths is given by (using q = 1 - p):

$$H(\text{geometric}) = -\sum_{j=0}^{\infty} q^j p \log(q^j p)$$
 (1)

and provides the optimal Shannon coding lower bound on the expected encoding length l per integer

$$E(l) \approx \sum_{j=0}^{\infty} q^{j} p\left(\lfloor j/m \rfloor + 1 + \lfloor \log m \rfloor\right)$$
 (2)

under the coin flip model. Thus, the Golomb code approaches the Shannon limit when  $q^m = 0.5$ . In particular, this ensures that for each integer j

$$-\log P(j) = \log(q^{j}p) \approx |j/m| + 1 + \lceil \log m \rceil \tag{3}$$

where P(j) is the probability associated with the integer j.

Finally, Golomb–Rice codes are a particularly convenient subfamily of Golomb codes, when  $m=2^k$  (Table 2). To encode j, we concatenate  $\lfloor j/2^k \rfloor$  1-bits, one 0-bit and the k least significant bits of j. The length of the encoding of j is thus  $\lceil j/2^k \rceil + k + 1$ . The decoding of Golomb–Rice codes is particularly simple, the position of the 0-bit gives the value of the prefix to be followed by the next k bits.

**Table 2.** Golomb–Rice encoding of integers j=0-33 with k=2 (m=4) and k=3 (m=8)

Number	Encoding $(k=2)$	Number	Encoding $(k=3)$	
0–3	0xx	0–7	0xxx	
4–7	10xx	8-15	10xxx	
8-11	110xx	16-31	110xxx	
33	11111111001	33	11110001	

Integer j is encoded by concatenating  $\lfloor j/2^k \rfloor$  1-bits, one 0-bit and the k least significant bits of j.

Table 3. Elias Gamma encoding

Number	Encoding
1	1
2–3	01x
4–7	001xx
8–15	0001xxx
16–31	00001xxxx

Each integer j is encoded by concatenating  $\lfloor \log j \rfloor$  0's with the binary value of j.

#### 3.2 Elias codes

In the Elias Gamma coding scheme, the preamble p(m) is a string of zeroes of length  $\lfloor \log j \rfloor$ , and the mantissa m(j) is the binary encoding of j. More precisely, to encode the scale and value of j:

- write  $|\log i|$  0-bits;
- followed by the binary value of j beginning with its most significant 1-bit.

The length of the encoding of j is  $2\lfloor \log j \rfloor + 1$  (Table 3). The decoding is obvious: first read n 0-bits until the first 1-bit is encountered, then read n more bits to get the binary representation of j.

Applying the relationship

$$-\log P(j) \approx 2|\log j| + 1 \tag{4}$$

to the integer probabilities, shows that Elias Gamma encoding asymptotically approaches the Shannon limit for  $P(j) \approx Cj^{-2}$ . This is a power–law relationship with exponent -2 and C is a normalizing constant. Note that for both Golomb [Equation (3)] and Elias Gamma codes [Equation (4)], several different consecutive integers can be encoded into a bit vector with the same length, hence the relationships  $-\log P(j) \approx \operatorname{length}(j)$  is only approximate with respect to geometric or power–law distributions over the integers. To be more precise, the optimal distribution associated with the Elias Gamma code can be separated into the product of a probability distribution over the length l given by  $P(l) = 2^{-l}$  and a uniform distribution over the integers having an encoding of length l given by  $P(j|l) = 2^{-l+1}$ .

More recently, new families of efficient fixed codes for integers have been developed (Baldi *et al.*, 2007; Hirschberg and Baldi, 2008; Moffat and Anh, 2006; Moffat and Stuiver, 2000), for instance, in the case of increasing or quasi increasing sequences of integers, by encoding only the deltas of the preambles. For sequence data, the absolute addresses are increasing, and the relative addresses could be made quasi-increasing if one were to apply a fixed permutation

to all the sequences to be stored, at the cost of storing and using this permutation (Baldi *et al.*, 2007).

# 3.3 Decoding and byte arithmetic

While the degree of compression achieved is an important criteria, the complexity and speed of decoding is also important in all the applications to be considered. For all the encoding algorithms described above, we have also described corresponding simple and fast decoding algorithms. Direct implementations of the decoding algorithms process the compressed representations bit-bybit; however, it is possible to implement even faster decoders, which decode the compressed data byte-by-byte. These faster decoders work by looking up information from precomputed tables. These tables are indexed by: (i) all possible bytes B (ranging from 0 to 255); and (ii) a bit-index i (ranging from 0 to 7) which marks the position of the decoder within the byte. These tables may store quantities such as the binary value of byte B starting from bit i, the number of bits turned on in byte B starting from bit i and the unary value of byte B starting from bit i. The exact quantities stored depend on the details of a particular decoder implementation. In practice, byte arithmetic considerably increases decoding speed, sometimes approaching as much as an 8-fold improvement over the corresponding bit-by-bit implementation. The exact value of the speedup depends on several factors including the characteristic of the data, the exact compression scheme and the hardware used.

#### 3.4 Variable codes

In genomic applications, in general the integers may not have a well-defined distribution, in which case it is always possible to use a general entropy encoding scheme, such as Huffman coding (Cover and Thomas, 1991; Huffman, 1952; McEliece, 1977), which essentially builds a prefix code by using a binary hierarchical clustering algorithm starting from the events (integers) with the lowest probability. While Huffman coding achieves compression close to the entropy limit, the price to pay over fixed coding schemes such as Golomb and Elias Gamma, or the more recent codes mentioned above, is the storage of the Huffman table which can be quite large in some applications. However, this is a fixed cost with respect to the database size, and therefore whether this cost is acceptable or not depends on the specific application. Small gains in compression over Huffman coding may be obtained using arithmetic coding (Rissanen and Langdonr, 1979; Witten et al., 1987), but at a non-trivial price in the complexity of computations.

# 4 RESULTS

# 4.1 Data extraction

To demonstrate the general approach, 3615 human mitochondrial sequences were downloaded from a recent version of GenBank. We focused on the sequences alone, ignoring any header and any other exogenous information. We first use the revised Cambridge Reference Sequence (rCRS) sequence (GenBank accession number: AC\_000021) as the reference sequence (Brandon *et al.*, 2005; Ruiz-Pesini *et al.*, 2007). The reference sequence is 16 568 bp long. Among the other sequences, 2671 correspond to complete genomes, while the remaining 944 correspond only to the coding region sequence, which is about ~1100 bp shorter than the full genome sequence, and extends from position 577 to 16 023 of the

reference sequence. Eighty sequences contained ambiguous symbols which, for simplicity, were replaced by the corresponding value in the reference sequence. This replacement is without much loss of generality since ambiguous symbols could easily be accommodated into the coding schemes, for instance as additional variation types.

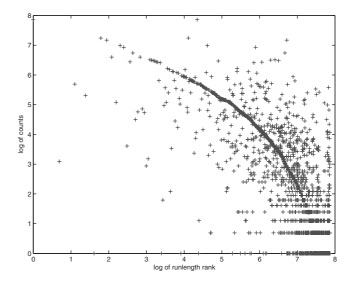
#### 4.2 General statistics

There are 4577 positions along the reference sequence where at least one of the other sequence deviates from the reference. In aggregate, there are 122 131 bp that deviate from the reference sequence. Besides substitutions, the total number of insertion and deletion events across all the sequences is 7119, the most frequent one being 1 bp insertions (4615 occurrences), followed by 2 bp deletions (901). Some well-known variants, such as the 'Asian-specific 9 bp deletion' (Harihara *et al.*, 1992; Thomas *et al.*, 1998), also occur frequently (255 occurrences). In total, there are 43 different kinds of variation events (Tables 6 and 7). On average, a given sequence deviates from the reference sequence in 33.8 bp with a SD of 13.43 bp. The average number of substitutions (transition/transversions) per sequence is 30.69 bp. The average number of insertions per sequence is 1.69 bp and the average number of deletions is 1.37 bp.

The distribution of the raw intervals using the rCRS as the reference sequence is represented in Figure 1 displaying the logarithm of the counts versus the logarithm of the rank (in decreasing order of frequency). Observed intervals vary from 0 to 14 997 bp, the most frequent one being an interval of 72 (2579 occurrences) (see interpretation in next section), followed by 687 (2418 occurrences), and followed by 5 (2130 occurrences). Overall this distribution is not strongly structured.

# 4.3 Changing the Reference Sequence

There are no particular reasons, beyond standardization and tradition, for using rCRS as the reference sequence. Furthermore, purely from a compression standpoint, the rCRS may not be optimal



**Fig. 1.** Distribution of intervals between variations using a log rank-log frequency plot. *x*-axis represents the logarithm of the rank associated with decreasing interval frequencies. *y*-axis represents the logarithm of the corresponding counts.

due to biases in data. To illustrate this point, we computed the haplotype distribution of the data using the simplified haplotype classification described in Figure 2 (see also, Brandon *et al.*, 2009; Mishmar *et al.*, 2003). We find the following skewed distribution: 11.2% African (405 sequences), 26.3% Asian (950 sequences) and 62.5% EurAsian (2260 sequences). In addition, it is well known that the original Cambridge Reference sequence contains a number of errors and has been revised over the years (Anderson *et al.*, 1981; Andrews *et al.*, 1999). (The revisions to the original sequence are described at: http://www.mitomap.org/mitoseq.html.) This alone, for instance, explains why the interval 72 is so frequent with respect to the rCRS: the rCRS sequence has a G in the corresponding position, which is a very rare variant, most likely an error.

Thus, it is clear that other reference sequences could be used to improve compression rates and minimize the total number of variants. Furthermore, the reference sequence does not need to be a sequence from an actual individual, but could be designed using purely statistical considerations. Note that the design of the reference sequence impacts not only the variants to be recorded, but also the intervals, and therefore it must also take into consideration any constraints a particular implementation may place on the intervals and their encodings. A reasonable choice adopted here to try to further improve the compression rate, is to use the consensus sequence, derived by computing the consensus at each position, as the reference sequence.

Using the consensus sequence, observed intervals vary from 0 to 11 717 bp, the most frequent one being an interval of 5 (2104 occurrences), followed by 1 (1251 occurrences) and followed by 259 (895 occurrences).

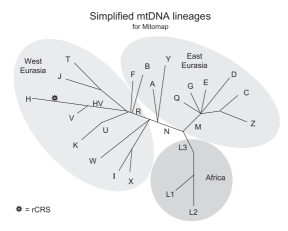


Fig. 2. Simplified haplotype classification used in Brandon et al. (2009).

# 4.4 Encoding and compression

We explored and compared different encoding schemes using both fixed and variable codes. The main sample of results is given in Tables 4 and 5 giving the average number of bits required to encode an interval or a variant, using Huffman, Golomb or Elias Gamma codes, with the rCRS or the consensus sequence, as well as the total number of bits required to encode the entire data. The Huffman coding tables for the events are given in Tables 6 and 7 for the rCRS and consensus sequence, respectively.

As can be seen in Table 5, Huffman coding achieves slightly better compression rates than Golomb or Elias Gamma coding, with a table storage cost that may be manageable in this case. The raw data takes 56 MB (58 817 584 bytes) of space. By concatenating the Huffman codes for the intervals and the variants (H+H), the encoded data requires only 167 KB of space, corresponding to a 345-fold level of compression. Using, for instance, Golomb codes for both the intervals and the variants (G+G) requires instead 195 KB. The choice of the reference sequence has a noticeable effect. Although the average number of bits required to encode an interval or a variant is slightly higher for the consensus sequence (Table 4), this is compensated by a considerable decrease in the total number of variants to be encoded. This is true here even with a consensus sequence that differs from the rCRS sequence by only 11 nt. As shown in Table 5, the same encoding method based on using two Huffman codes (H+H), applied with the consensus sequence, requires only 133 KB to store the entire data. This corresponds to a 433-fold level of compression, roughly a 23% improvement.

## 5 DISCUSSION

A simple but general data structure and data encoding approach has been developed for the efficient storage of genomic data. The approach specifically leverages homology between sequences and is different from general compression algorithms for text, or compression algorithms for single genome data (Behzadi and

**Table 5.** Total file size comparison using the rRCS and the consensus sequence, with Huffman encoding for both intervals and variants (H+H), or Golomb encoding for both interval and variants (G+G), or Elias Gamma encoding for both interval and variants (E+E)

H+H	G+G	E+E
167 (345)	195 (295)	226 (254)
133 (433)	159 (361)	183 (314)
	167 (345)	167 (345) 195 (295)

Numbers are given in Kilobytes (1024 × 8 bits). In comparison, the raw data takes 56 MB (57439.05 KB). Compression factor are given in parenthesis.

Table 4. Comparison of the average bit cost of encoding intervals and events for Huffman, Golomb and Elias Gamma encoding schemes using the rCRS and the consensus sequence

	Intervals	Intervals			Variants		
	Huffman	Golomb	Elias Gamma	Huffman	Golomb	Elias Gamma	
Cambridge	9.21	11.10	12.93	2.66	2.44	2.77	
Consensus	9.75	12.03	13.86	2.44	2.59	2.97	

Table 6. Huffman encoding for the event types using the rCRS

Table 7. Huffman encoding for the event types using the consensus sequence

Variant	Count	Binary code	Variant	Count	Binary code
G	42839	11	C	26164	11
C	24753	01	A	19576	01
T	22345	00	G	18002	00
A	21003	101	T	16528	101
InsC	3980	1001	InsC	3980	1001
Del2bp	901	100011	Del2bp	901	100011
Del1bp	757	100001	Del1bp	757	100001
InsCC	360	1000100	InsCC	360	1000100
InsT	313	1000001	InsT	313	1000001
Del9bp	255	1000000	Del9bp	255	1000000
InsA	222	10001011	InsA	222	10001011
InsCCC	34	1000101000	InsCCC	34	1000101000
InsCCCC	30	10001010110	InsCCCC	30	10001010110
InsG	29	10001010100	InsG	29	10001010100
InsCCCCC	16	100010101110	InsCCCCC	16	100010101110
InsACA	15	100010101011	InsACA	15	100010101011
InsCCCCCC	12	100010100110	InsCCCCCC	12	100010100110
InsCCCCCCC	8	1000101010101	InsCCCCCCC	8	1000101010101
InsAC	6	1000101001011	InsAC	6	1000101001011
InsCCT	5	1000101001010	InsCCT	5	1000101001010
Del6bp	4	100010101111100	Del6bp	4	100010101111100
Del8bp	4	10001010111101	Del8bp	4	10001010111101
InsCCCCCTCTA	3	10001010011111	InsCCCCCTCTA	3	10001010011111
Del3bp	3	10001010101001	Del3bp	3	10001010101001
InsGC	3	10001010011101	InsGC	3	10001010011101
InsCCCCCCC	3	10001010101000	InsCCCCCCC	3	10001010101000
InsTT	3	10001010011110	InsTT	3	10001010011110
Del4bp	3	10001010011100	Del4bp	3	10001010011100
InsACAC	2	1000101011111100	InsACAC	2	1000101011111100
InsACACA	1	100010100100000	InsACACA	1	100010100100000
InsCCCCCCCC	1	100010100100111	InsCCCCCCCC	1	100010100100111
InsTA	1	10001010111111110	InsTA	1	10001010111111110
InsGA	1	10001010111111101	InsGA	1	10001010111111101
InsGG	1	10001010111111100	InsGG	1	10001010111111100
InsCA	1	1000101011111011	InsCA	1	1000101011111011
InsAG	1	10001010111111010	InsAG	1	10001010111111010
InsGATCACAG	1	100010100100011	InsGATCACAG	1	100010100100011
Del10bp	1	100010100100010	Del10bp	1	100010100100010
InsTCTCTGTTCTTTCAT	1	100010100100001	InsTCTCTGTTCTTTCAT	1	100010100100001
InsACACAC	1	100010100100101	InsACACAC	1	100010100100101
InsAGAA	1	100010100100100	InsAGAA	1	100010100100100
InsCACA	1	1000101011111111	InsCACA	1	1000101011111111
Del5bp	1	100010100100110	Del5bp	1	100010100100110

Deletions (Del) are followed by their length. Insertions (Ins) by their content.

Deletions (Del) are followed by their length. Insertions (Ins) by their content.

Fessant, 2005; Chen *et al.*, 2002; Williams and Zobel, 1997). The approach has been demonstrated on the mitochondrial genomes, where it leads to 2–3 orders of magnitude improvement in data storage. From these compact representations, full sequences can be recovered rapidly using the reference sequence. Furthermore, queries regarding the existence and nature of variants at particular coordinate positions, such as those arising in a variety of applications from medicine to forensics, can be answered efficiently. Additional encryption methods may be applied to these representations to protect the security of both the genomic data and the queries.

The approach has been used for lossless compression, however it could be used also in lossy compression, for instance, by ignoring variants that are not medically relevant. The approach is also

applicable to other kinds of sequences, such as RNA or protein sequences. While for demonstration purposes we have used a single reference sequence, it is clear that one could cluster the data and use different reference sequences for different subgroups. In the case of mitochondria genomes, for instance, Figure 2 would suggest using at least three different reference sequences. Whether the gain in compression that can be expected for each subgroup, akin to the gain achieved by going from the rCRS to the consensus sequence, is worth the cost of having multiple reference sequences rather than a single one, cannot be answered in generality and depends on the details of a particular application, the number of genomes to be stored coming from each group and so forth. For future work, the same idea of multiple reference sequences can be extended beyond

Table 8. Compression of the read addresses information from three HTS experiments (see text)

Encoding	DataSet 1	DataSet 2	DataSet 3
Raw Sequence	133 366 560	353 182 128	8 869 613 600
Flat File	75 525 168	185 536 864	8 396 646 344
Elias Gamma absolute	358 402 (210.73)	79 281 140 (2.34)	1 373 892 116 (6.11)
Elias Gamma relative	185 542 (407.05)	27 741 238 ( <b>6.69</b> )	340 764 564 ( <b>24.64</b> )
Monotone Value (MOV)	169 664 ( <b>445.15</b> )	39 528 754 (4.69)	834 672 652 (10.06)

The size in bits for the raw sequence data, the corresponding flat text file format for the corresponding addresses, and the compressed files for different compression algorithms. Elias Gamma coding is applied both to the absolute and relative addresses. Compression factors with respect to the flat text file format are given in parentheses, with top compression factors in bold. MOV is a coding algorithm specifically designed for increasing sequences of integers described in Baldi *et al.* (2007).

the storage of genomes within a given species, to the storage of genomes from multiple species by using a phylogenetic hierarchy of reference sequences.

Finally, the approach can be extended to human nuclear genomes (see also, Christley et al., 2009) and to HTS from different technologies and different kinds of experiments. For human SNP variation, data and statistics are readily available (Goldstein and Cavalleri, 2005; Hinds et al., 2005; The International HapMap Consortium, 2007). A comprehensive list of human SNPs is available from the dbSNP database maintained by NCBI. The current release (version 129) contains about 15 million SNPs. This data can readily be compressed using the techniques described here and additional gains in compression can be achieved by storing separately a fixed table recording the location of all the SNPs and leveraging the skewed distribution of some of the SNP variants. In preliminary experiments, we have achieved compression factors of over 1000 on the raw HapMap sequence data. Although SNPs account for most of genetic variation events between individuals, a much larger fraction of the genome (in terms of the total number of bases) is involved in larger structural variation events, such as copy number variations (CNVs). While there have been studies attempting to derive a preliminary assessment of large-scale genomic complexity and variation (Feschotte and Pritham, 2007; Tuzun et al., 2005), statistics on the frequencies and location of these more complex structural variations in the human genome are still at an earlier stage of development. For instance, comparative analysis of the single diploid genome described in Levy et al. (2007), 'revealed more than 4.1 million DNA variants, encompassing 12.3 Mb. These variants (of which 1 288 319 were novel) included 3 213 401 single nucleotide polymorphisms (SNPs), 53 823 block substitutions (2206 bp), 292 102 heterozygous insertion/deletion events (indels) (1571 bp), 559 473 homozygous indels (182 711 bp), 90 inversions, as well as numerous segmental duplications and copy number variation regions. Non-SNP DNA variation accounts for 22% of all events identified in the donor, however they involve 74% of all variant bases. This suggests an important role for non-SNP genetic alterations in defining the diploid genome structure'. A better statistical understanding of the coding constraints posed by these complex events, and how to encode them, should become possible as more full human genome sequences become available in the coming years (www.1000genomes.org).

Regarding HTS data, for illustration purposes, here we consider the problem of storing the genomic addresses of the reads from three HTS datasets associated with different HTS technologies. The first dataset is obtained from the laboratory of Dr S. Sandmeyer at UCI and comes from an experiment aimed at mapping retrotransposon Ty3 insertion sites in the yeast genome. It consists of 833 541 sequence reads, all of length 19 bp. The second dataset comes from a chromatin immunoprecipitation assay (ChIP-Seq) used to map the in vivo binding site locations of the neuron-restrictive silencer factor (NRSF) in humans (Johnson et al., 2007). It consists of 1 697 991 sequence reads, all of length 25 bp and mapped to the most recent human genome sequence (hg18). The third dataset corresponds to a full diploid human genome sequencing experiment for an Asian individual (Wang et al., 2008). This is a very large dataset with enough reads to provide 36-fold average coverage, and we utilize the existing mapping of the reads provided by the YH database (Li et al., 2009) to the human reference genome. For illustrative purposes, we report only the results corresponding to the reads associated with chromosome 22. For chromosome 22, there are 31 118 532 reads that vary in length from 30 to 40 bp for a total of 1 108 701 700 bp of sequence data. While complete details of these experiments will be reported elsewhere, Table 8 shows the resulting compression factors which are again in the range of 1-3 orders of magnitude, depending on the statistical properties of the datasets. The same techniques described here can readily be applied to storing also the length of the reads, the content of the reads, where they differ from the reference genome, their quality and so forth. Statistical properties of the reads and the underlying HTS technologies, e.g. increasing error rates towards the end of the read, can also be exploited to achieve efficient compression. Thus, the data structures and compression algorithms described here provide a framework for the management of HTS and genomic data that can be flexibly applied in different environments.

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